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**A2**

(54) Title: RECOMBINANT INFLUENZA A VIRUSES

(57) Abstract: The invention relates to a recombinant NS gene of an influenza A virus comprising a functional RNA binding domain and a gene sequence modification after nucleotide position 400 of the NS1 gene segment, counted on the basis of influenza A/PR/8/34 Virus, wherein the modification bars transcription of the remaining portion of the NS1 gene segment. It further relates to embodiments, wherein the modification comprises deletions, insertions, or a shift of the open reading frame, and particularly to constructs comprising an insertion of an autocleavage site 2A, the nef gene from HIV-1 or the sequence encoding the ELDKWA-epitope of gp41 of HIV-1. The invention also relates to influenza virus transfectants that contain the modified NS gene and have an IFN inducing phenotype but which may or may not be sensitive towards IFN. The invention also relates to vaccines comprising such a chimeric virus.

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RECOMBINANT INFLUENZA A VIRUSES

TECHNICAL FIELD

The invention is in the fields of vaccine development and application
5 and relates to attenuated live vaccine vectors, more specifically to such
vectors based on or derived from genetically modified influenza A virus
strains, and to the manufacture of recombinant influenza viruses and vaccines.

BACKGROUND OF THE INVENTION

10 Influenza viruses are segmented negative-strand RNA viruses and
belong to the *Orthomyxoviridae* family. Influenza A virus consists of 9
structural proteins and codes additionally for one nonstructural NS1 protein
with regulatory functions. The non-structural NS1 protein is synthesized in
large quantities during the reproduction cycle and is localized in the cytosol
15 and nucleus of the infected cells. The segmented nature of the viral genome
allows the mechanism of genetic reassortment (exchange of genome
segments) to take place during mixed infection of a cell with different viral
strains. Several features make influenza viruses attractive candidates for the
development of effective live vaccine vectors against different diseases:
20 (i) influenza viruses induce strong cellular and humoral immune responses, at
the systemic and the mucosal level against viral proteins following infection;
(ii) influenza virus as an RNA virus does not contain a DNA phase in its
replication cycle. Therefore chromosomal integration of viral genes into the
host can be excluded;
25 (iii) many different influenza virus subtypes are available. Since antibodies
against these different subtypes show no or little crossreactivity, pre-existing
immunity to the viral vector in the host, which is frequently a problem for
other live vectors, can be circumvented. Also, effective booster immunizations
with different subtype influenza viruses expressing the same antigens might
30 be possible; and
(iv) attenuated influenza viruses as live influenza vaccines, which were shown
to be safe and immunogenic in humans, are available.

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Until now the main problem of utilizing influenza virus as a vector concerns the size of the virus genome and its limited capacity to tolerate foreign sequences. Among ten influenza viral proteins, only the surface glycoproteins haemagglutinin (HA) and neuraminidase (NA) have been 5 successfully engineered for stable expression of foreign epitopes. Since influenza virus tolerates an insertion of only approximately 10 amino acids into its HA molecule, there is only a limited possibility to influence the conformation properties of inserted epitopes which would probably be better presented if longer sequences would be introduced. Besides, surface influenza 10 glycoproteins such as HA or NA cannot be considered as optimal targets for the presentation of foreign sequences because of their association with antigenic properties of the viruses. An HA live virus construct containing a desired foreign antigen is not applicable for boosting immunizations (e.g., by second and further administrations) because of the pre-existing immunity 15 against the HA caused by the first immunization or by a natural virus infection. A booster immunization would be possible only upon introduction of the desired antigenic structure into another HA molecule belonging to a different influenza virus subtype. It is evident that such a process is difficult, laborious and extremely time consuming and therefore unlikely to be suitable 20 for routine vaccine preparation

Preceding investigations in connection with the present invention have indicated that the NS gene of influenza A virus may be a promising alternative to HA as a viral carrier for presenting a desired foreign antigen to the animal or human immune system. The recently established method of reverse 25 genetics (*Egorov et al.*, 1998, J Virol 72(8), 6437-41) allows to rescue influenza viruses containing long deletions or insertions of foreign sequences at the carboxyl side of the non-structural Protein 1 (NS1 protein). NS1 protein is abundant in influenza virus-infected cells and stimulates cytotoxic T- lymphocyte (CTL) responses as well as antibody responses during the natural 30 course of influenza virus infection.

Further details about the influenza virus NS gene can be found in WO 99/64571. Additionally, WO 99/64571 discloses that attenuated influenza A

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virus transfectants containing knockout deletions of the entire NS1 gene were found to have a strong interferon (IFN) inducing phenotype. This was concluded from the finding that such transfectants were able to grow on IFN deficient Vero cells but were unable to grow on hen eggs or Madin-Darby

5 Canine Kidney (MDCK) cells.

The influenza NS1 protein is an RNA-binding protein which has been implicated in a number of regulatory functions during influenza virus infection. It is synthesized in large amounts and found mainly in the nucleus early during infection and later in the viral cycle in the cytoplasm of the infected

10 cells. Other than the influenza NS-1 protein, another regulatory viral protein, namely the Nef protein of HIV-1 which is a myristylated protein, is localized in the cytosol in association with the cell membrane.

An immune response directed against early expressed regulatory HIV-1 proteins could possibly allow the elimination of virus-infected host cells in the 15 replication cycle before release of new infectious viral particles would even occur. As the Nef protein is among the first ones to be released and further is one of the major HIV proteins produced following infection, it could play a crucial role in developing an efficacious anti-AIDS vaccine.

The HIV-1 "negative factor" (Nef) is encoded by an open reading frame 20 which is located at the 3' end of the virus, partially overlapping the U3 region of the 3' long terminal repeat. Up to 80% of the early, multiply spliced class of viral transcripts encode Nef. The Nef gene product is an NH₂-terminally myristylated protein of 27 to 30 kDa, which is predominantly localized in the cytoplasm and associated with the membrane and the cytoskeletal matrix. It 25 is well conserved among the different human (HIV-1 and HIV -2) and Simian immunodeficiency viruses (SIV).

The close evolutionary relationship between these primate lentiviruses suggests that the Nef protein plays an important role in viral infection and pathogenesis, although the exact role in the virus life cycle and its functions 30 at the cellular level are still the subject of current research.

Various details about the Nef protein and its effects are already known, however. For instance, it is reported that some humans infected with Nef-

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deleted HIV remained disease-free, with normal CD4 counts 10 to 14 years after infection, although deletion of Nef is not a universal finding in long-term nonprogressors. In addition, Nef-deficient SIV fails to produce AIDS in infected adult macaques. SIV mutants deleted for the Nef gene even induce protection
5 against a virulent challenge. Nef was shown to stimulate HIV-1 proviral DNA synthesis and its expression has also been found to induce the efficient internalization and degradation of the cell surface CD4 receptor for HIV-1. This Nef-induced CD4 down-regulation, which renders cells resistant to viral superinfection, has the potential to increase virus replication by facilitating
10 release of progeny virions. It was further demonstrated that extracellular Nef protein could activate HIV-1 from latent to productive infection both in infected T-cell lines and in PBMC from asymptomatic carriers. Further, it was shown that CTLs inefficiently lysed primary cells infected with HIV-1, if the viral Nef gene product was expressed.

15 Protection of HIV-infected cells from efficient recognition and killing by CTLs correlates with the Nef-mediated down-regulation of MHC class I molecules. Nef also interferes with the induction of IL-2 mRNA in T-cell lines. Furthermore, there are a large number of cellular partners that have been found to be associated with Nef expression including Src family kinases, β -
20 COP, a serine-threonine kinase, thioesterase and p53.

It is also reported that the majority (about 2/3) of HIV-1 seropositive patients generated Nef-specific CTLs.

Two central multirestricted immunodominant regions (amino acids 66 to 100 and 115 to 146) and a carboxyl-terminal region (amino acids 182 to 206)
25 were identified within the Nef protein. These three multirestricted immunodominant regions (amino acid sequences containing more than one T-cell epitope) are being recognized by human CD8+ CTLs in association with at least 14 different MHC class I molecules including the important MHC haplotypes HLA-A1, -A3, -A11, -B8, -B17, -B18 and -B37.

30 The two central multirestricted domains of the Nef protein are the most highly conserved regions among different HIV-1 isolates and were immunodominant for most of the asymptomatic HIV-1 seropositive donors

tested. In addition to the high immunogenicity of the HIV-1 Nef protein, which has been demonstrated by its capacity to induce strong T-cell immune responses, also Nef-specific B-cell immune responses are reported in the literature.

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SUMMARY OF THE INVENTION

The present invention relates to the development of attenuated live vaccine vectors, more specifically to such vectors based on or derived from genetically modified influenza A virus strains. It further relates to the

- 10 construction and modification of genetically engineered non-structural genes of influenza A viruses, particularly of the NS1 gene segment, wherein the modifications include deletions of selected parts of the NS1 gene segment and/or insertions of heterologous, preferably antigenic, sequences into selected sites of the NS1 gene. It is another objective of the invention to
- 15 provide chimeric influenza viruses containing such modified NS1 gene segments but which do not suffer from the drawback of being IFN sensitive, in contrast to the transfectants disclosed in WO 99/64571. The invention further relates to recombinant proteins obtained from the NS1-modified viruses by expression in a suitable host system and further to a vaccine
- 20 comprising the NS1-modified viruses of the present invention.

- It is yet another objective of the invention to provide a method for obtaining recombinant influenza viruses as well as attenuated influenza vaccines, based on the generation of continuous cell lines (e.g. Vero, MDCK etc.) expressing synthetic influenza genes (minus sense RNA) comprising natural or
25 engineered influenza sequences (deletions or insertions). These cell lines producing high quantities of such genes can be used for infection with influenza virus followed by selection procedures in order to get a gene of interest incorporated into the viral progeny.

- The present inventors have established a reverse genetics system on
30 Vero cells allowing them to manipulate the virulence of the PR8 influenza A virus strain by changing the length of the translated NS1 protein. In the course of the research leading to the present invention the capacity of

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influenza A virus to tolerate and present long insertions in the NS gene have been investigated. A collection of several chimeric NS1 gene constructs using heterologous sequences including the HIV-1 derived sequences encoding ELDKWA of gp41 or Nef, has been established by insertion of one or more of 5 the heterologous sequences or, optionally, several repeats of any such sequence, in frame into the NS1 protein.

The aforementioned heterologous sequences were inserted downstream nt position 400 (corresponding to aa position 124) and optionally preceded by the 2A autocleavage site sequence and/or by a leader sequence derived from 10 the influenza HA molecule. Other constructs additionally comprised an anchor sequence derived from the influenza HA molecule as an insertion right after the desired antigenic sequence(s), thus forming the end of the entire heterologous insertion. In each case the insertions were followed by a stop codon to prevent transcription and translation of the remaining portion of the 15 NS1 gene segment (including the effector domain), while maintaining the cleavage site for the NS splicing (necessary for transcription and translation of the NS2 = NEP gene segment) fully functional.

Rescued viruses caused expression and accumulation of the foreign antigens in the cytosol and/or on the surface of the infected cells. The 20 inventors also successfully rescued transfected viruses harbouring a multirestricted immunodominant region rich in T-cell epitopes of HIV-1 Nef protein (136 amino acids).

All transfecteds displayed normal growth characteristics in Vero cells, embryonated chicken eggs and MDCK cells, but were attenuated in mice. 25 Chimeric influenza NS1-Nef viruses did not replicate in respiratory tracts of infected mice, but were able to induce a strong Nef-specific CTL response following a single intranasal immunization. In addition, a Nef-specific antibody response was detected following three immunizations. Transfer of the recombinant NS-nef gene by genetic reassortment from the viral PR 30 (influenza A/PR/8/34; Egorov et al., 1994, Vopr. Virusol. 39:201-205) vector to other influenza strains resulted in the same level of attenuation and immunogenicity. This finding permitted the present inventors to perform

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effective boosting immunizations using several attenuated vectors of different antigenic subtypes.

Thus, the inventors were able to demonstrate that the approach to create a set of influenza chimeric strains belonging to different influenza 5 subtypes while bearing the identical recombinant chimeric NS1 gene, gives the opportunity to create several strains for boosting immunizations. They have further proven that once a new chimeric NS1 gene construct is rescued it can be routinely transferred to another influenza strain by genetic reassortment.

10 DESCRIPTION OF THE FIGURES

Fig.1 shows a functional map of the engineered NS1 protein of PR8;

Fig. 2 shows the structure of recombinant NS1 proteins of the rescued influenza transfectant viruses expressing gp-41 and IL-1 β peptides;

15 Fig. 3 shows the structure of recombinant NS1 protein of the rescued influenza transfectant PR82Anef (PR8/Nef) expressing aa70-206 of the Nef protein of HIV-1 NL4-3;

Fig. 4 shows the replication of chimeric influenza/Nef viruses in mouse lower respiratory tracts;

20 Fig. 5 shows the number of IFN-gamma secreting cells from immunized mice per 10^6 spleen cells detected after immune spleen cells were incubated in the presence of Nef peptide, NP peptide or without peptide as an indicator for T-cell responses;

Fig. 6 shows the number of IFN-gamma secreting cells from the lymph 25 nodes draining the respiratory tracts from immunized mice per 10^6 spleen cells detected after immune spleen cells were incubated in the presence of Nef peptide, NP peptide or without peptide as an indicator for T-cell responses;

Fig. 7 shows Nef-specific serum IgG immune responses following 2nd 30 and 3rd Immunizations of mice immunized with recombinant influenza/Nef viruses;

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Fig. 8 exhibits results of a plaque reduction assay;
Fig. 9 exhibits results of an interferon induction assay;
Fig. 10a-10c show immunofluorescence of Vero cells infected
previously with recombinant PR8/Nef virus (10a, 10b) and the wild type
5 influenza PR8 virus (10c).

Fig. 11 shows Nef peptide-specific and NP peptide-specific immune
responses as a quantification of IFN- γ secreting cells in murine spleen cells.

Fig. 12 shows Nef peptide-specific and NP peptide-specific immune
responses as a quantification of IFN- γ secreting cells in lymph nodes draining the
10 respiratory tracts of immunized mice.

Fig. 13 shows Nef peptide-specific and NP peptide-specific immune
responses as a quantification of IFN- γ secreting cells in urogenital single cell
populations of immunized mice.

Fig. 14 exhibits results of an ELISA assay determining Nef-specific IgG in
15 sera of mice two weeks after the third immunization with the PR8/NS-Nef or the
Aichi/NS-Nef vector.

Fig. 15 is a schematic representation of an Influenza NS transcription
system for expression of minus sense RNA for use in generating recombinant
influenza viruses.
20

DETAILED DESCRIPTION OF THE INVENTION

In one embodiment, the invention relates to genetically engineered NS
gene constructs of an influenza A virus comprising sequence modifications,
25 i.e. deletions or insertions, between nucleotide (nt) positions 400 and 525 of
the NS1 gene segment (numbering is based on the NS gene of influenza
A/PR/8/34 virus). Unexpectedly, it turned out that maintaining functionality of
the NS1 gene segment up to nt position 400 (corresponding to aa position
124 of the NS1 protein) while concomitantly deleting the remaining portion or
30 at least a major part thereof or inserting a foreign nt sequence into the region
after nt position 400 or shifting the reading frame to cause wrong
transcription and translation of the remaining NS1 portion resulted in the

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rescue of NS gene constructs that rendered their viral vectors IFN inducing but not IFN sensitive.

This surprising finding was confirmed by experiments wherein the chimeric influenza viruses engineered according to the present invention not 5 only induced a strong IFN response in MDCK cells and hen eggs but were also able to grow on these host substrates at an efficiency comparable to the wildtype PR8 virus. In contrast, chimeric influenza viruses containing deletions of the first third of the NS1 gene or deletions of the entire NS1 gene displayed an IFN inducing as well as an IFN sensitive phenotype. They were 10 unable to grow on hen eggs or MDCK cells and therefore could only be cultivated on IFN deficient cell lines such as Vero cells. Chimeric influenza viruses of the latter type have been disclosed in WO 99/64571. It is assumed that the viruses of the present invention, which are not as strongly attenuated as the viruses of WO 99/64571, are more immunogenic and therefore better 15 suitable for the manufacture of highly effective live vaccines against various kinds of viral infections.

In another embodiment of the invention the genetically engineered NS gene is used as a genomic fragment of influenza A virus in a method wherein it is transferred to any desired influenza A virus strains or live influenza 20 vaccines by means of genetic reassortment. In this context, it is preferred that the genetically engineered NS gene is established as a cDNA clone that can be transferred to any influenza A virus strain or live influenza vaccine as a genomic fragment by means of reverse genetics methods. For example, another vector such as Aichi/NS-Nef belonging to the H3N2 subtype, but 25 containing the same recombinant NS-gene, can be obtained in this manner. In contrast to strategies generating recombinant influenza viruses expressing foreign antigens in the context of HA or NA molecules, this approach enables a fast generation of a set of non-crossreactive vectors for optimal boosting immunizations.

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In a preferred embodiment of the invention the genetically engineered NS gene is designed for the expression of viral antigens, particularly for expression of the HIV-1 sequences of Nef or ELDKWA of gp-41.

In another embodiment of the invention the genetically engineered NS 5 gene is rescued as a genomic fragment of influenza virus expression of which contributes as or elicits a factor of protein kinase p-68 (PKR) overexpression and activation in infected cells.

In another embodiment of the invention the chimeric NS gene is part of an attenuated (cold adapted) live influenza vaccine vector wherein the 10 genetically engineered NS gene is the main factor or an additional factor of attenuation. It is particularly useful for the manufacture of safe and highly effective, influenza virus-based vaccines including but not restricted to anti-HIV-1 vaccines, wherein the transfected chimeric NS gene construct comprises gene sequences of nef, 2A, and/or gp-41 or other viral antigens, 15 for the induction of strong antibody and/or B- and T-cell immune responses.

The vaccine comprising an attenuated (cold adapted) live influenza virus vector can be prepared in a suitable pharmaceutical formulation and may be used for prophylactic immunizations as well as for therapeutic vaccination, including induction of IFN release in combination with a stimulation of B- and T- 20 cell response. In such formulations influenza vectors might be used in combination with any other vector expressing analogous antigens to ensure a maximum booster effect. Thus, generating of attenuated influenza NS vectors offers the possibility to obtain novel recombinant vaccines with nearly optimal balance of safety and immunogenicity directed against a broad range of 25 pathogens.

In a particular embodiment of the invention the genetically engineered NS gene of an influenza A virus comprises a heterologous nucleotide insertion derived from the HIV-1 nef gene (nucleotides 210-618 of the nef gene of the HIV-1 clone NL4-3) plus an insertion of the autocleavage sequence 2A (54 30 nucleotides) N-terminally to the HIV-1 derived insertion at the position 400 of the NS protein. Elimination of the first 68 amino acids of the Nef protein was done to exclude domains comprising the myristylation site and other domains

associated with pathogenic properties of the multifunctional HIV-1 Nef protein.

In further experiments the inventors found that influenza PR8/Nef virus and influenza Aichi/Nef virus resulted in a high titer of antibodies against the viral vector and a less but still significant titer against the nef gene. The

5 PR8/Nef virus is a PR8-124 virus with truncated NS1 containing a 2A autocleavage sequence after aa position 124 of the NS1 protein which additionally comprises a nef sequence (aa 70-206 of HIV-1 Nef protein) following the autocleavage site. The Aichi virus is reassorted in that except for the NS gene all genes including the genes encoding the envelope proteins
10 HA and NA originate from H3N2 Aichi wild type virus, while the recombinant NS gene originates from the PR8/Nef virus. It was also observed that the PR8/Nef and Aichi/Nef viruses caused strong T-cell responses against the nef gene as well as the viral vector. This experiment proved that it is possible to transfer the chimeric NS gene into another influenza virus strain to elicit
15 essentially the same immune response. This finding is important as it allows to provide for possibilities to boost immunizations and to design seasonal influenza vaccines with varying immunogenic subtypes but constant chimeric NS1 gene-based activity.

In another embodiment the invention provides for a method for
20 generating recombinant influenza viruses by constructing a vector comprising a modified NS gene wherein the NS1 gene sequence is partially or entirely deleted or truncated, mixing said vector with lipids to allow self-assembling of lipid-DNA complexes and transfecting the lipid-DNA complexes into a desired continuous cell line, for example a Vero or MDCK cell line, and selecting
25 clones that stably integrate and replicate the modified NS gene, which then are infected with any desired influenza strain, and particularly, with an epidemic wild-type influenza strain, to produce attenuated viral progeny containing said modified NS gene. In this method, the modified NS gene may further comprise insertions of heterologous gene sequences coding, for
30 instance, for other viral antigens or pathogens e.g. such as the ones disclosed herein.

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- It is another object of the present invention to provide a method for rapid vaccine manufacture comprising the steps of transforming a continuous cell line to produce a desired synthetic viral gene, particularly a modified NS gene of influenza A virus wherein the NS1 gene is partially or entirely deleted 5 or truncated, infecting the transformed cell line with a desired virus, particularly, with an epidemic wild-type influenza strain, to produce attenuated viral progeny containing said modified NS gene, selecting attenuated recombinant viruses and multiplying such viruses under conditions suitable for efficient virus replication, preferably using interferon-deficient substrates, and 10 combining harvested virus material with a pharmaceutically acceptable carrier resulting in an anti-viral vaccine. In this method, the modified NS gene may further comprise insertions of heterologous gene sequences coding, for instance, for other viral antigens or pathogens e.g. such as the ones disclosed herein.
- 15 Further embodiments are defined in the dependent claims. In order that the invention described herein may be more fully understood, the following examples are set forth. The examples are for illustrative purposes only and are not to be construed as limiting this invention in any respect.

20 Example 1: Preparation of recombinant negative strand influenza A viruses
(“reverse genetics method”)

The plasmid clones containing the nef sequence have been prepared on the basis of the existing plasmid clone of influenza NS gene pUC19/NSPR (*Egorov et al.*, 1998, J Virol 72(8), 6437-41). The nef sequence was inserted into the 25 NS1 protein ORF, downstream of an additional sequence: a protease recognition sequence P2A (NFDLLKLAGDVESNLG/P) derived from foot and mouth disease virus that is posttranslationally cleaved by an ubiquitous cellular protease (*Mattion et al.*, 1996, J Virol 70(11), 8124-7; *Percy et al.*, 1994, J Virol 68(7), 4486-92), so that the gp-41 molecule should be cleaved 30 from the NS1 polypeptide and transported to the cell surface. The plasmid clone was used for synthesis of chimeric RNA to be transfected into Vero cells in order to rescue the recombinant influenza viruses. In the functional

- map of the engineered NS1 protein of the present invention (Fig. 1) it is indicated that the insertions are introduced after aa position 124 and followed by a stop codon which has in effect that the remaining adjacent portion of the NS1 gene segment (including the effector domain) rests untranslated. From 5 Fig. 2 it can be understood how the desired antigenic or otherwise heterologous sequences (e.g. gp-41 and IL-1 β sequences) may be arranged to yield immunogenic constructs that after transfection into a suitable viral vector, preferably a cold adapted influenza virus, could form the basis of an effective vaccine against various infectious diseases.
- 10 Analogously, Fig. 3 shows the arrangement of insertion of aa70-206 of the Nef protein HIV-1 NL4-3 into the NS1 protein of the rescued influenza transfectant PR82Anef(PR8/Nef).

In general, the experiments showed a tendency wherein the length of the heterologous insert or inserts was directly proportional to the degree of 15 attenuation of the resulting virus strain. Additionally, the immunogenic potential of the expression products of larger inserts usually exceeded the one of smaller inserts. Therefore, it is preferred according to the present invention to make inserts encoding at least about 80 amino acids.

To create the chimeric Aichi/NS-Nef virus the RNA representing the 20 recombinant NS segment of the PR8/NS-Nef virus was introduced into the genome of influenza A/Aichi/1/68 (H3N2) virus by a standard genetic reassortment performed on Vero cells utilizing rabbit polyclonal anti PR8 virus hyperimmune serum for selection. Genotyping of reassortants was performed by RT-PCR amplification and comparative restriction analysis of cDNA copies 25 derived from each genome segment.

Example 2: Transfection of recombinant viruses in Vero cells

Synthetic negative sense RNA have been derived from plasmid clones by T3 transcription in the presence of purified viral RNP. Vero cells were 30 previously infected with the helper influenza virus reassortant strain 25A-1 (H1N1, (Egorov et al., 1994, Vopr Virusol 39(5), 201-5) and then transfected

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with RNA complexes by DEAE-dextran transfection (*Egorov et al.*, 1998, J Virol 72(8), 6437-41; *Luytjes et al.*, 1989, Cell 59(6), 1107-13). Rescued transfectant viruses have been plaqued, purified on Vero cells 3 times, amplified on Vero cells and checked for biological properties.

- 5 Fig. 10a-c show immunofluorescence of Vero cells infected previously with recombinant PR8/Nef virus (MOI 0,01 in 10a and 0,1 in 10b) and the wild type influenza PR8 virus (10c). 24 hrs following infection cells were trypsinized and fixed on cover slides with 100% acetone. After several wash steps in PBS the slides were incubated for 40 min at 37°C with a 1:50
10 dilution of anti Nef (aa179-195 epitope) mouse monoclonal antibody, then washed twice in PBS and incubated with 1:100 dilution of goat anti mouse IgG FITC conjugated antibody.

Example 3: Immunization of BALB/c mice

- 15 Groups of three BALB/c mice each were immunized with $2\text{-}5 \times 10^5$ PFU /mouse of influenza viruses PR8/Nef, Aichi/Nef; PR8-124 and with 4×10^4 PFU/mouse of the PR8wt as indicated in Fig. 5. Spleen cells from immunized mice were obtained 9 days later and used as effector cells in the ELISPOT assay. Figure 5 shows the number of IFN-gamma secreting cells per 10^6
20 spleen cells detected after immune spleen cells were incubated in the presence of the EWRFDSRLAFHHVAREL peptide (Nef peptide), TYQRTRALVRTMGD peptide (NP peptide) or without peptide (w/o peptide). Results are expressed as average +/- SEM of duplicate cultures.

- Groups of three BALB/c mice were immunized with $2\text{-}5 \times 10^5$ PFU/
25 mouse of influenza virus PR8/Nef, Aichi/Nef; PR8-124 and 2×10^4 PFU/mouse of the PR8wt as indicated in Figure 6. Simple cells suspensions from the lymph nodes draining the respiratory tracts from immunized mice were obtained 9 days later and used as effector cells in the ELISPOT assay (Power et al, J Immunol Methods 227:99-107): Briefly, threefold serial dilutions of
30 cell populations derived from murine spleens, draining lymph nodes and the urogenital tracts were transferred to wells coated with anti IFN- γ mAb (R4-

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6A2; BD PharMingen). Cells were incubated for 22 hours at 37°C and 5% CO₂ in DMEM medium containing 10% FCS, IL-2 (30 U/ml), penicillin, streptomycin and 50µM 2-ME in the presence of synthetic peptides. A biotinylated anti IFN-γ mAb (XMG1.2; BD PharMingen) was utilized as a conjugate antibody, then plates were incubated with streptavidin peroxidase (0.25 U/ml; Boehringer Mannheim Biochemica). Spots representing IFN-γ secreting CD8⁺ cells were developed utilizing the substrate 3-amino-9-ethylcarbazole (Sigma) containing hydrogen peroxide in 0.1 M sodium acetate, pH 5.0. The spots were counted with the help of a dissecting microscope and results were expressed as the mean number of IFN-γ secreting cells ± SEM of triplicate cultures. Cells incubated in the absence of synthetic peptides developed < 10 spots/10⁶ cells. Since depletion of CD8⁺ cells resulted usually in > 92% reduction of spot formation, cell separation was omitted in most assays. Figure 6 shows the number of IFN-gamma secreting cells per 10⁶ spleen cells detected after immune spleen cells were incubated in the presence of the EWRFDSRLAFHHVAREL peptide (Nef peptide), TYQRTRALVRTMGD peptide (NP peptide) or without peptide (w/o peptide).

From Fig. 4, which shows the replication of the chimeric influenza/Nef viruses in mouse lower respiratory tracts, it can be understood that the PR8/Nef and Aichi/Nef viruses did not replicate in that tissue, hence were strongly attenuated, while at the same they were highly immunogenic to the mice causing strong T-cell and B-cell immune responses (as shown in Figures 5, 6 and 7). To characterize the insert (Nef peptide)-specific and vector (NP-peptide)-specific CD8⁺ T cell response female BALB/c mice were immunized once or twice i.n. without narcosis with 10⁶ PFU per animal of the PR8/NS-Nef, Aichi/NS-Nef; PR8/NS-124 or PR8 w.t. virus.

Three BALB/c mice per group were immunized once or twice i.n. in the absence of anesthesia with 10⁶ PFU/mouse of influenza PR8/NS-Nef, Aichi/NS-Nef, PR8/NS-124 or PR8 w.t. as indicated in Fig. 11. The booster immunization was performed 21 days after priming. The single cell suspensions obtained 10 days after immunization from spleens of mice were assessed for Nef peptide-

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specific (A) or NP peptide-specific (B) IFN- γ secreting CD8 $^{+}$ T cells in an ELISPOT assay. Fig. 11 shows the mean numbers of antigen-specific IFN- γ secreting cells \pm SEM of triplicate cultures.

Lymph nodes draining the respiratory tracts (mediastinal and retrobronchial 5 lymph nodes) were collected 10 days after immunization from immunized BALB/c mice as described in the Fig. 11. Single cell suspensions were assessed for Nef peptide-specific (A) or NP peptide-specific (B) IFN- γ secreting CD8 $^{+}$ T cells in an ELISPOT assay. Fig. 12 shows the mean numbers of antigen-specific IFN γ -secreting cells \pm SEM of triplicate cultures.

10 Three BALB/c mice per group were immunized twice i.n. in the absence of anesthesia with 10⁶ PFU/mouse of influenza PR8/NS-Nef, Aichi/NS-Nef or PR8/NS-124 virus as indicated in Fig. 13. The booster immunization was performed 21 days after priming. The single cell suspensions derived 10 days after the second immunization from digested urogenital tracts (vagina, cervix, 15 uterine horns and urethras) of immunized mice were assessed for Nef peptide-specific (A) or NP peptide-specific (B) IFN- γ secreting CD8 $^{+}$ T cells in an ELISPOT assay. Fig. 13 shows the mean numbers of antigen-specific IFN γ secreting cells \pm SEM of triplicate cultures.

Mice immunized once with either the PR8/NS-Nef or Aichi/NS-Nef virus 20 induced significant numbers of Nef peptide-specific CD8 $^{+}$ T cells in single cell suspensions derived from spleens (139 \pm 4 spots in PR8/NS-Nef immunized mice; 137 \pm 39 spots in Aichi/NS-Nef immunized mice; Fig. 11B) and lymph nodes (173 \pm 23 spots in PR8/NS-Nef immunized mice; 160 \pm 25 spots in Aichi/NS-Nef immunized mice; Fig. 12B). No relevant Nef peptide-specific CD8 $^{+}$ 25 T cell response was determined in both compartments of mice immunized with the PR8/NS-124 and PR8 w.t. viruses (the number of spots were always lower than 13; Fig 11B and 12B).

When vector (NP peptide)-specific CD8 $^{+}$ T cell responses were 30 compared, similar numbers of specific CD8 $^{+}$ spleen cells were found in all groups of mice tested (Fig. 11A). In contrast to the systemic compartment (spleens), significant differences were obtained in the mucosa-associated

respiratory lymph nodes. The replication competent PR8/NS-124 virus induced a markedly higher frequency of NP-peptide specific CD8⁺ T cells. whereas recombinant influenza/NS-Nef viruses and the pathogenic PR8 w.t. virus induced lower magnitudes of NP peptide-specific CD8⁺ T cells (Fig. 12A).

5

In Fig. 7, which displays Nef-specific serum IgG immune responses of mice immunized with recombinant influenza/Nef viruses, the B-cell immune responses following 2nd and 3rd immunizations have been demonstrated to be strongest in the case where the mice have been immunized twice by PR8/Nef 10 followed by a third immunization with Aichi/Nef, while the response was less prominent with twice immunizations using the PR8-124 virus.

Further, a group of mice was primed i.n. with the PR8/NS-Nef virus and boosted 21 days later with the Aichi/NS-Nef virus. Another group of mice was immunized with the same viruses but in the reverse order. Data shown in Fig. 15 11 and 12 indicate that the sequence in which the respective recombinant vectors were used for priming and boosting appeared to be crucial, since it was consistently observed that priming with the Aichi/NS-Nef (H3N2) followed by boosting with the PR8/NS-Nef (H1N1) induced a significantly lower number (approximately the range of the primary CD8⁺ T cell response) of the Nef 20 peptide-specific and NP peptide-specific CD8⁺ T cells in spleens and draining lymph nodes when compared with the reverse order of immunization. A strong secondary antigen specific CD8⁺ T cell response was detected in both of the compartments tested after priming the mice with the recombinant PR8/NS-Nef (H1N1) vector followed by a boost using the H3N2 subtype Aichi/NS-Nef 25 vector. In this case, Nef- and NP peptide-specific secondary responses were approximately 1.5 to 3 times higher than after a single immunization (Fig. 11 and 12).

Single cell suspensions derived from the urogenital tracts were obtained from immunized mice. Two immunizations were necessary before significant 30 numbers of Nef peptide-specific CD8⁺ T cells could be detected. The strongest Nef peptide specific CD8⁺ T cell response was detected when mice were

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primed i.n. with PR8/NS-Nef (H1N1) virus and subsequently boosted with the Aichi/NS-Nef (H3N2) virus (342 ± 18 IFN- γ SC/ 10^6 cells; Fig. 13B). This immunization protocol was also found to induce the strongest NP peptide-specific CD8 $^+$ T cell response (Fig. 13A).

5 As described for the detection of T-cell responses mice were utilized to assess the Nef-specific serum antibody response. Mice were primed i.n. either with 10^6 PFU/ml of the PR8/NS-Nef (H1N1) or Aichi/NS-Nef (H3N2) virus and were boosted three weeks later with the same vector. The third immunization was performed following three more weeks utilizing the vector of the different
10 subtype. The control group was immunized three times with the PR8 w.t. virus. The reactivities of serum samples (obtained two weeks after the third immunization) with the GST-Nef fusion peptide were determined by ELISA and are shown in Fig. 14. Nef-specific antibodies were detected only in groups of mice which had been successively immunized with H1N1 and H3N2 vectors (Fig.
15 14). The highest level of Nef-specific IgG was detected in mice immunized twice i.n. with 10^6 PFU of the PR8/NS-Nef (H1N1) virus and boosted with 10^6 PFU of the Aichi/NS-Nef (H3N2) virus as compared with the control group which had been immunized three times i.n. with PR8 w.t. virus (Fig. 14).

Both influenza virus vectors (PR8/NS-Nef and Aichi/NS-Nef) were
20 completely attenuated in mice, since no viral titers could be detected in mouse respiratory tissues. These attenuated phenotypes of both recombinant viruses indicate that introduction of additional amino acids downstream of the position 125 of the NS1 protein can affect some function of the NS1 protein since PR8/NS-124 virus encoding the same size of the NS1 protein grew efficiently in
25 mouse respiratory tracts (Fig. 4). The low efficiency of the 2A site to cleave Nef antigen from the N-terminal part of NS1 protein, especially at the late stage of infection, might be responsible for additional attenuation, although a direct effect of the Nef polypeptide interacting with some intracellular components can not be excluded.

30 The data indicate that completely attenuated recombinant influenza/NS-Nef viruses are capable to induce a primary CD8 $^+$ T cell response directed to

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the inserted Nef polypeptide in spleens and in lymph nodes draining the respiratory tracts of mice immunized i.n. without anesthesia. At the same time, vector (NP peptide)-specific CD8⁺ T cell responses in spleens and draining lymph nodes of mice immunized i.n. either with the PR8/NS-Nef or 5 Aichi/NS-Nef vector were in the range of those induced by the virulent PR8 w.t. virus although the strongest NP-peptide specific CD8⁺ T cell response detected in the draining lymph nodes was found in mice immunized with the PR8/NS-124 virus efficiently replicating in the lungs.

The results indicate that it is possible to achieve a similar effect utilizing 10 influenza vectors belonging to different antigenic subtypes. Importantly, influenza virus vectors are capable to induce a CD8⁺ T cell response circumventing the pre-existing immunity caused by a different influenza virus subtype.

The immunogenic potential of attenuated influenza/NS-Nef vectors might 15 be explained by the fact that viruses containing truncated forms of the NS1 protein induce high levels of type 1 interferons in vivo. Nef-expressing vectors as well as PR8/NS-124 virus induced significantly higher levels of type 1 interferons in serum following immunization of mice if compared with the corresponding w.t. parent viruses (data not shown).

20

Example 4: Plaque reduction assay

MDCK cells were treated for 24 h with a supernatant from MDCK cells infected by delNS virus (same construct as disclosed in WO 99/64571, i.e., containing entire NS1 deletion) as a known potent IFN alfa/beta inducer (Fig. 25 8). The content of the IFN alfa/beta was estimated to 100 U following an overnight treatment with pH2. The results in Fig. 8 are given in log of plaque forming units (PFU reflecting the differences in viral titers on IFN alfa/beta treated and untreated cells.

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Example 5: Interferon induction

MDCK cells were infected for 24 h with 5 MOI of different influenza viruses as outlined in Fig. 9. The supernatants were treated overnight with pH 2 at 4°C for virus inactivation. Treated supernatants were adjusted to pH 7.4 with 1N NaOH. Twofold serial dilutions of these supernatants were added to MDCK monolayers for 24 hrs and 50 PFU of the vesicular stomatitis virus (VSV) were then added per well. The results represent the dilution of the supernatant at which VSV plaque formation was reduced by 50%.

10 Example 6: A method for accelerated production of recombinant viruses and anti-viral vaccines.

a) Generating recombinant cell lines producing modified NS gene:

A plasmid vector was constructed according to the schematic representation in 15 Fig. 15. The NS gene was cloned into the backbone of a pS65T-C1 vector (Clontech), using the CMV promotor to initiate transcription. Inserting NS in reverse orientation (3' end towards CMV promotor) leads to the transcription of minus sense RNA. Transcription is terminated by a hepatitis delta virus (HDV) sequence that comprises a self-cleaving RNA site. Hence, this vector contains 20 an influenza A NS gene where the cassette of the multiple stop codons is introduced at nt position 140 in a manner such that translation of this gene (reading frame 3) leads to a truncated form of the NS1 protein (comprising only 38 amino acids). It was found that influenza A viruses expressing such a short NS1 protein are highly attenuated in animals (Egorov et al. 1998, J Virol. 72, 25 8, p 6473).

Sequence of HDV (85nt):

TGGCCGGCATGGTCCCAGCCTCGCTGGCGCCGGCTGGCAACATTCCGAG
GGGACCGTCCCCCTCGGTAAATGGCGAATGGGAC

Sequence of PR8NS38: (906 nt)

AGCAAAAGCAGGGTGACAAAGACATAATGGATCCAAACACTGTGTCAAGCTT
TCAGGTAGATTGCTTCTTGCATGTCGCCAACCGAGTTGCAGACCAAGAAC
5 TAGGTGATGCCCATTCCTGATCGGCTCGCCAGTGAATAACTAGCTGAAT
CAGAAATCCCTAAGAGGAAGGGGCAGCACCCCTCGCTGGACATCGAGACAG
CCACACGTGCTGAAAGCAGATAGTGGAGCGGATTCTGAAAGAAGAACATCGA
TGAGGCACTTAAATGACCATGCCCTGTACCTGCGTGCCTACCTAACCTG
ACATGACTCTGAGGAAATGCAAGGGACTGGTCCATGCTATACCCAAGCAG
10 AAAGTGGCAGGCCCTTTGATCAGAATGGACCAGGGATCATGGATTAAGA
ACATCATACTGAAAGCGAACTTCAGTGTGAATTGACCCGCTGGAGACTCTA
ATATTGCTAAGGGCTTACCGAAGAGGGAGCAATTGTTGGCGAAATTCA
ATTGCCCTCTCCAGGACATCTGCTGAGGATGTCAAATGAGTGGAG
TCCTCATGGGGGACTTGAATGGAATGATAACACAGTTGAGTCTGAAACT
15 CTACAGAGATTGCTTGGAGAACAGTAATGAGAATGGAGACCTCA
CTCCAAAACAGAAACGAGAAATGGCGGGACAATTAGGTCAAGAGTTGAAG
AAATAAGATGTTGATTGAAGAAGTGGAGACACAAACTGAAGATAACAGAGAA
TAGTTTGAGCAAATAACATTATGCAAGCCTTACATCTATTGCTGAAGTGG
GCAAGAGATAAGAACATTCTCGTTCAGCTTATTAGTAATAAAAACACCC
20 TGTTCTACT

This plasmid vector was used for transfection in order to transform Vero cells. Before transfection, cells were seeded in appropriate cell culture flasks (e.g. 25 cm) and incubated at 37°C until having reached about 50 % confluence. To improve the transfection efficiency the cationic lipid reagents (lipofectin, lipofectamine 2000) may be preincubated in medium without serum prior to mixing it with the plasmid-DNA. Accordingly, 2-6 µl cationic lipid reagent were diluted in 100 µl OPTI-MEM I serumfree transfection medium and incubated 30 min at room temperature. Meanwhile 1-2 µg DNA were diluted in 100 µl OPTI-MEM I, then mixed with the lipid solution and incubated 15 min at room temperature. While DNA-lipid complexes formed, the cells were washed twice with serum-free transfection medium to remove residual proteins. The transfection cocktail was diluted to a total volume of 1 ml using transfection medium and added to the cells. Cells were incubated at 37°C with 7 % CO₂ from 5 to 8 h. Thereafter cell culture supernatant was removed and cells were fed with normal culture medium. 24 hours post transfection stable transfecants were selected by addition of selection medium containing genetin sulfate G418 (400 µg/ml). Three weeks later

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stable transfectants appeared which were subcloned by limiting dilutions. Several subclones were tested for their ability to rescue the ts helper virus (25A-1 mutant, which is a reassortant virus wherein the NS gene responsible for the ts phenotype originates from cold-adapted influenza strain 5 A/Leningrad/134/47/57, and the remaining genes originate from PR8 virus; Egorov et al., 1994, Vopr. Virusol.39: 201-205) at a temperature of 40° C. For that purpose, the cells were infected with the 25A-1 mutant and incubated at 40° C for 72 hours. Those subclones yielding viral progeny that contained influenza viruses carrying the recombinant NS gene were selected 10 and further multiplied under conditions allowing for efficient cell growth.

b) Recombinant influenza virus and vaccine production:

Transformed cells expressing a modified NS gene segment comprising a truncated NS1 gene with or without insertions, preferably as described 15 hereinbefore, are infected with a desired influenza virus, particularly with a wildtype epidemic virus, under conditions as described above and incubated to allow for the development of viral progeny wherein the wildtype NS gene is replaced by the recombinant modified NS gene supplied by the transformed host cells. The viral yield can be cloned by plaqueing methods (e.g. negative 20 colonies under agar overlay) on a normal Vero cell line and each viral colony can be screened for the presence of the modified recombinant NS gene by the RT-PCR method and/or by other methods (depending on how the gene was modified). Positive viral plaques are then purified by further plaqueing purification steps. Finally, the recombinant influenza strains are highly 25 attenuated because of the truncated NS protein. They serve as vaccine candidates and are multiplied preferably using IFN deficient substrates such as Vero cells, young chicken embryos (less than 10 days old) and the like, for rapid manufacture of highly attenuated live influenza vaccines.

It is pointed out, however, that the method described in this Example is 30 just one way to exemplify the underlying general concept of preparing and establishing mammalian cell lines, particularly immortalized or continuous cell

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lines, that after transformation with any desired viral gene sequences stably integrate and express these sequences, and thus allow for relatively simple and rapid design and manufacture of reassorted, recombinant viruses of whatever origin.

ABBREVIATIONS

aa	amino acid
CMV	cytomegalovirus
CTL	cytotoxic T-lymphocyte
5 gp	glycoprotein
HA	haemagglutinin
IFN	interferon
Ig	immunoglobulin
IL	interleukin
10 i.n.	intranasally
mAb	monoclonal antibody
MHC	major histocompatibility complex
NA	neuramidase
nef	negative factor of HIV
15 NP	nucleoprotein
NS	non-structural
ORF	open reading frame
RNP	ribonucleoprotein
SIV	Simian immunodeficiency virus
20 ts	temperature sensitive
v	viral
VSV	vesicular stomatitis virus
wt	wild type

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Claims

We claim:

1. A recombinant NS gene of an influenza A virus comprising a functional RNA binding domain and a gene sequence modification after nucleotide 5 position 400 of the NS1 gene segment, counted on the basis of influenza A/PR/8/34 virus, wherein the modification bars transcription of the remaining portion of the NS1 gene segment.
2. The recombinant NS gene according to claim 1, wherein the modification comprises deletions, insertions, or a shift of the open reading frame.
- 10 3. The recombinant NS gene according to claim 1, wherein the modification comprises an insertion of at least one sequence selected from the group consisting of an autocleavage site 2A, the nef gene from HIV-1, the sequences encoding the ELDKWA or ELDKWAS epitopes of gp41 of HIV-1, the sequence encoding IL-1 β or a part thereof, a leader sequence, and an 15 anchor sequence.
4. The recombinant NS gene according to claim 3, wherein insertion encodes a sequence of at least 80 amino acids.
5. A genetically engineered, preferably cold adapted, influenza virus comprising a modified NS gene as defined in any one of claims 1 to 4.
- 20 6. The influenza virus according to claim 5, having an IFN inducing phenotype.
7. The influenza virus according to claim 5 or 6, which is not IFN sensitive.
8. A vaccine comprising at least one virus, preferably a mixture of different viruses, as defined in any one of claims 5 to 7, in a suitable pharmaceutical 25 formulation.
9. The vaccine according to claim 8, for prophylactic or therapeutic application against a viral infection, preferably against influenza or HIV-1 infection.

10. A method for the manufacture of recombinant influenza viruses comprising the steps of:

transforming a mammalian cell, preferably a continuous cell line, with a DNA vector comprising a modified NS gene segment wherein the NS1 5 gene sequence is partially or entirely deleted or truncated, selecting transformed cells that express the modified NS gene segment, infecting the selected cells with a desired influenza virus, preferably a wildtype epidemic strain, incubating the infected cells to allow for the development of viral progeny containing the modified NS gene segment, and selecting 10 and harvesting said viral progeny containing the modified NS gene segment.

11. The method according to claim 10, wherein transformation of the mammalian cell is accomplished comprising mixing said vector with lipids to allow for a self-assembly of the lipid and the DNA to form lipid-DNA

15 complexes, and incubating the mammalian cells in the presence of said lipid-DNA complexes resulting in an uptake of the complexes into the cells, and preferably into the cell nucleus.

12. The method according to claim 10 or 11, wherein the DNA vector is a transcription system for minus sense influenza RNA.

20 13. The method according to any one of claims 10 to 12, wherein said viral progeny containing the modified NS gene segment is further combined with a pharmaceutically acceptable carrier for use as an attenuated influenza live vaccine.

14. A live attenuated influenza virus vaccine comprising at least one 25 genetically engineered, preferably cold adapted, influenza virus as defined in claim 5, preferably a mixture thereof, obtainable in a method according to any one of claims 10 to 13.

Fig. 1

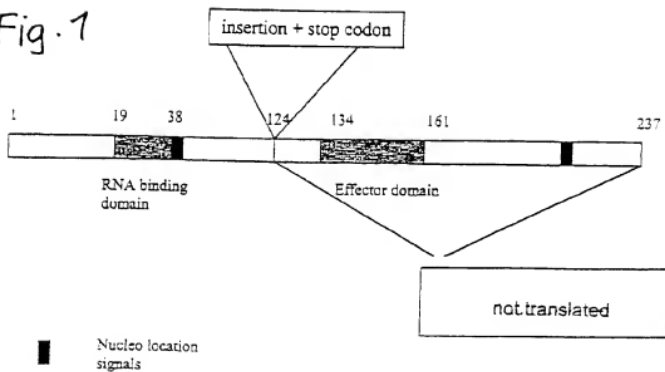


Fig. 2

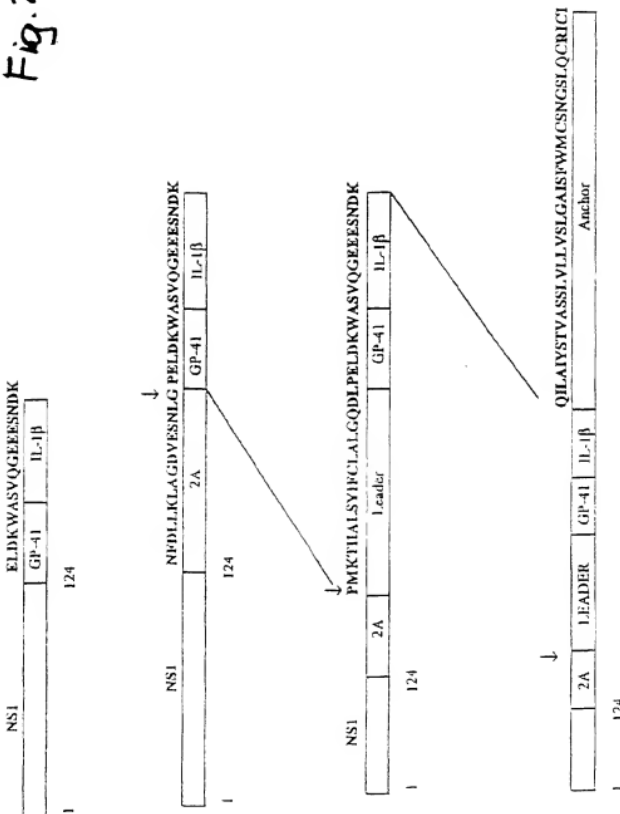


Fig. 3

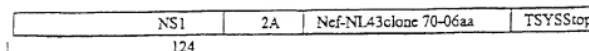
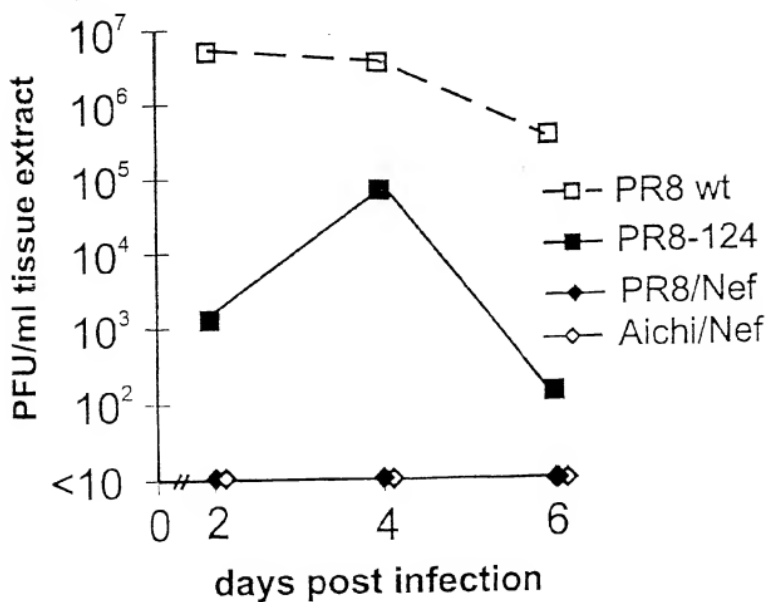


Fig. 4



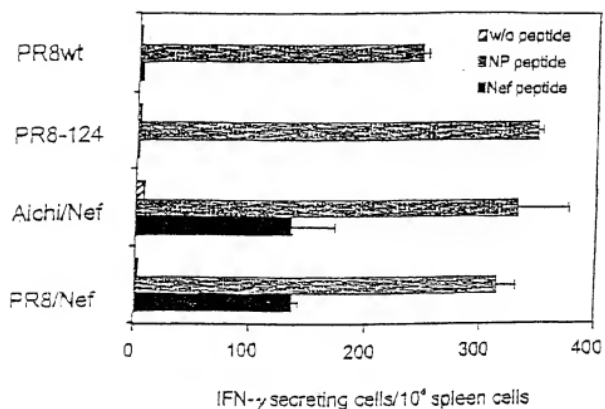


Fig. 5

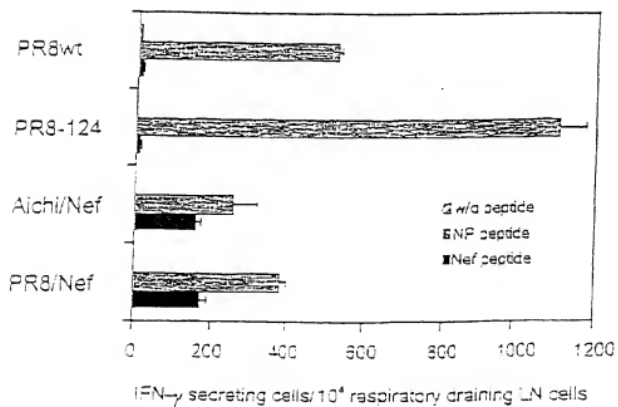


Fig. 6

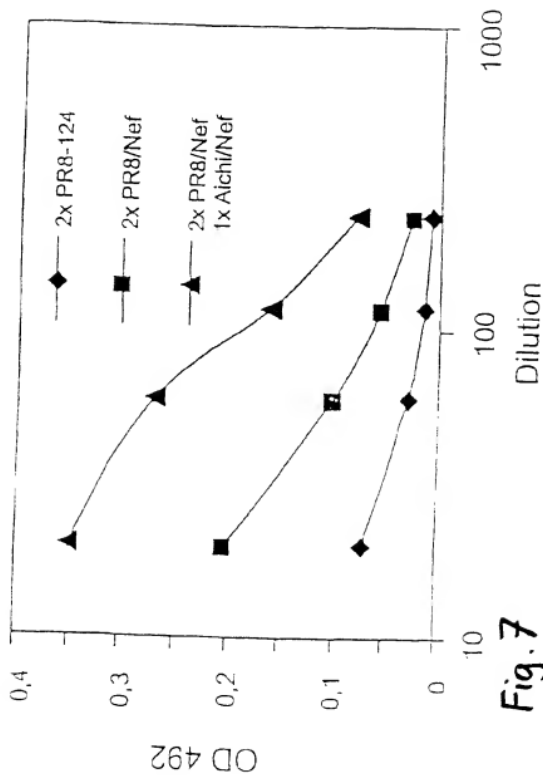


Fig. 7

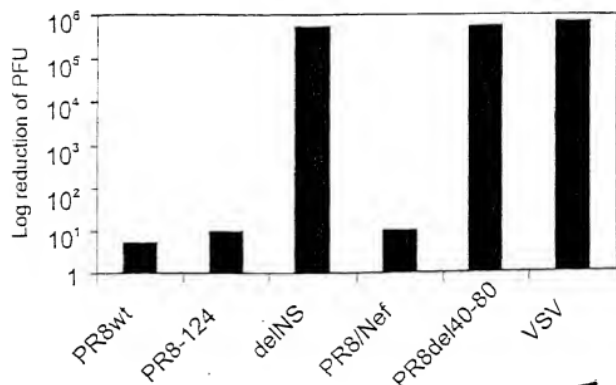


Fig. 8

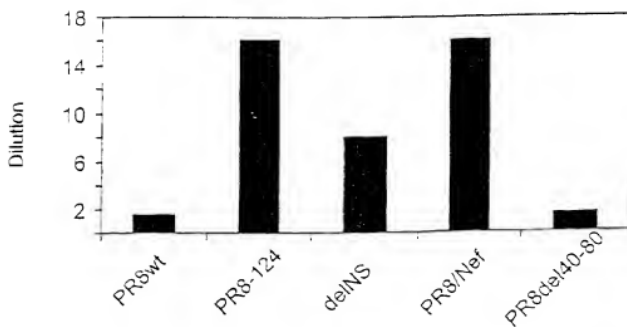


Fig. 9

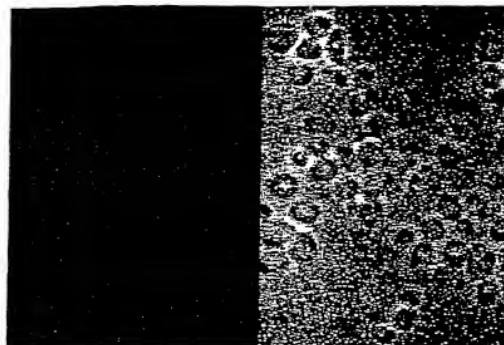


Fig. 10a

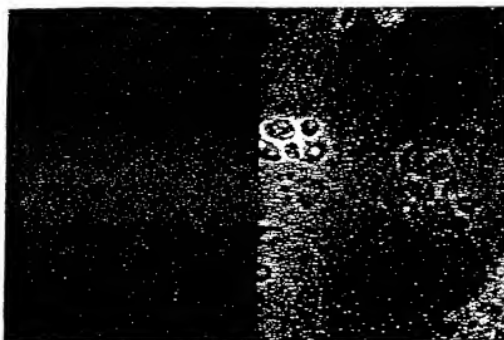


Fig. 10b

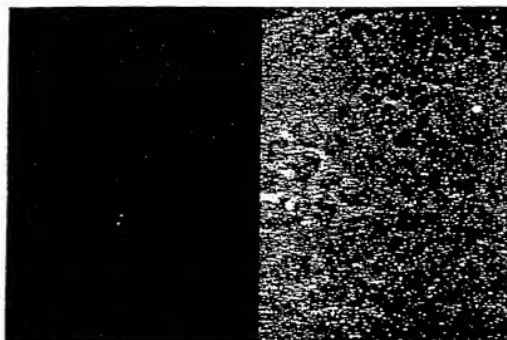
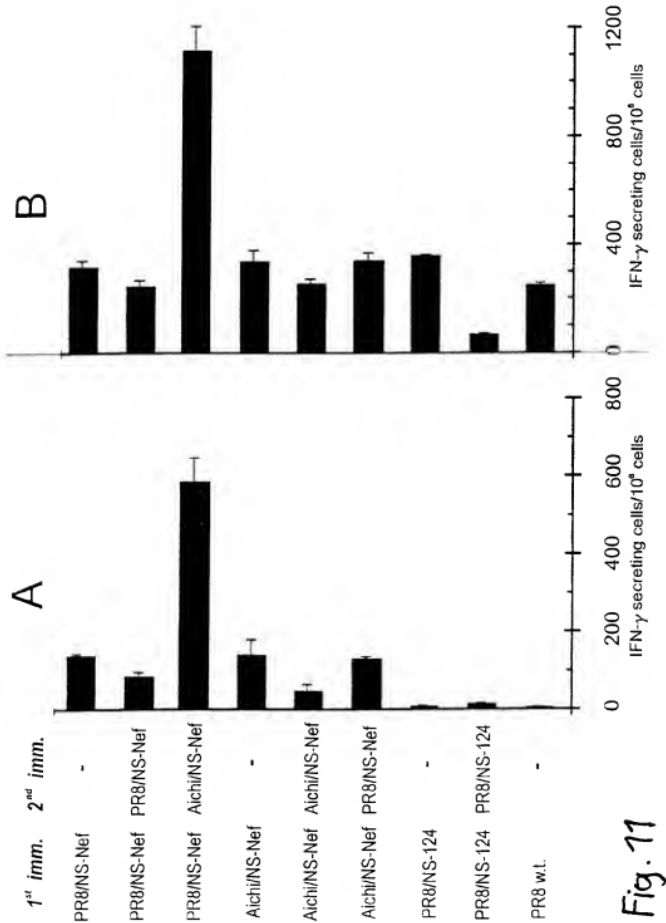
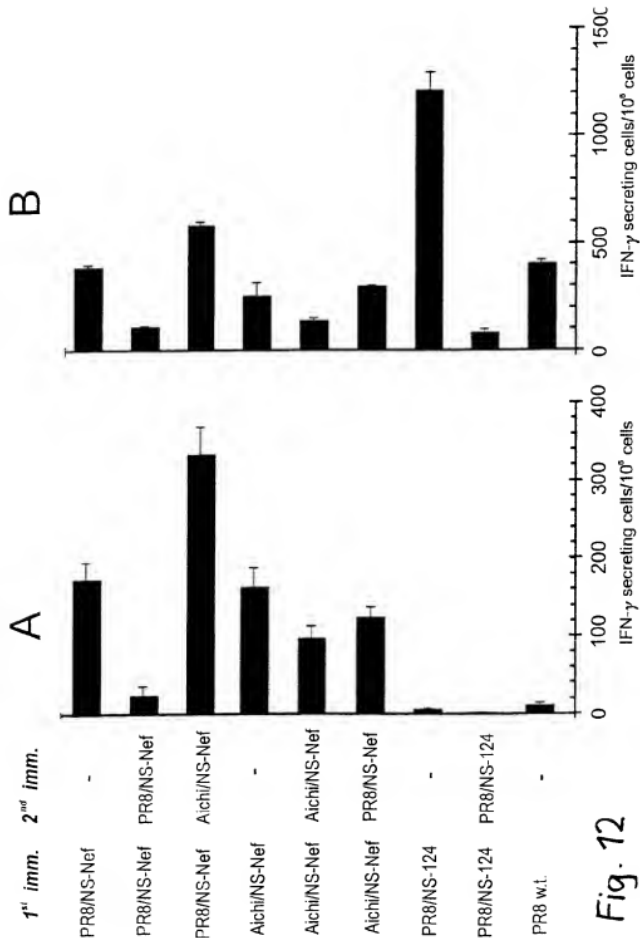


Fig. 10c

**Fig. 11**

**Fig. 12**

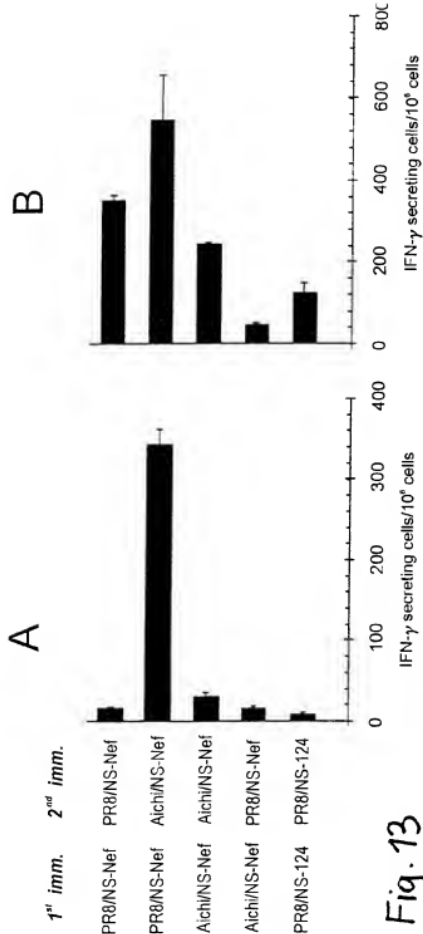
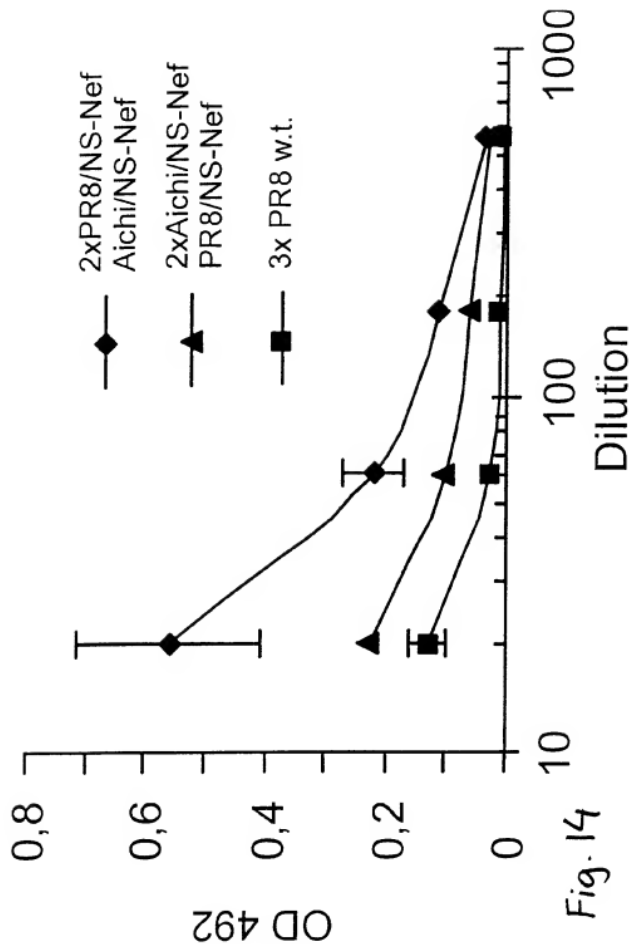


Fig. 13



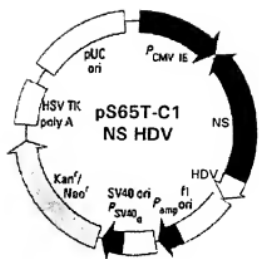


Fig. 15